

Plant Morphology of Heterotrimeric G Protein Mutants

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The heterotrimeric G protein complex, comprising $G\alpha$, $G\gamma$ and $G\gamma$ subunits, is an evolutionarily conserved signaling molecular machine that transmits signals from transmembrane receptors to downstream target proteins. Plants conserved the core G protein elements, while developing their own regulatory systems differently from animals. Genetic evidence supports the conclusion that the heterotrimeric G proteins regulate shoot, root and epidermis development, as well as sugar sensing, hormone responsiveness and abiotic and biotic stress tolerance. This review is a compendium of the known morphological changes conferred by loss- and gain-of-function mutations of the G protein subunit genes across three higher land plant models, namely Arabidopsis, rice and maize.

Keywords: AGB1 • GPA1 • CT2 • d1 • DEP1 • GS3.

Abbreviations: AGB1, Arabidopsis G protein β subunit 1; AGG,Arabidopsis G protein γ subunit; CT2, Compact Plant2; DEP1, Dense and Erect Panicle 1; d1, dwarf1; GCR1, G protein-coupled receptor 1; GPA1, G protein α subunit 1; GPCR, G protein-coupled receptor; GS3, Grain Size 3; IM, inflorescence meristem; QTL, quantitative trait locus; RGS, regulator of G protein signaling; SAM, shoot apical meristem; 7TM, seven transmembrane; XLG, extra-large G protein.

Introduction to G Protein Signaling

Animal heterotrimeric G proteins serve as physical couplers between seven transmembrane (7TM) G protein-coupled receptors (GPCRs) and downstream components designated as effectors (Kaziro et al. 1991). G proteins have three subunits: $G\alpha$, $G\beta$ and $G\gamma$, among which the $G\alpha$ subunit binds a guanine nucleotide: GDP or GTP. A ligand-bound GPCR induces exchange of GDP for GTP on $G\alpha$ leading to its conformational change and G protein complex dissociation. The active $G\alpha$ or $G\beta\gamma$ subunits then interact with downstream effectors and modulate their activities. Intrinsic GTP hydrolysis by $G\alpha$ returns it to the GDP-bound, basal state. Regulator of G protein signaling (RGS) proteins accelerate GTP hydrolysis by $G\alpha$, thereby suppressing G protein activity. Plants lack the conventional G protein regulation by GPCRs, because their G proteins spontaneously activate themselves without GPCRs (Johnston et al.

2007, Urano et al. 2012). Plants have G protein-coupled receptor 1 (GCR1), a 7TM protein weakly homologous to the Dictyostelium cAMP receptor (Colucci et al. 2002); however, its action on G proteins remains equivocal (Chen et al. 2004, Pandey et al. 2006). Most vascular plants, except cereals, utilize a 7TM RGS protein to modulate their G protein activity (Chen et al. 2003, Urano et al. 2012), although the entire regulatory system still remains unclear (Urano et al. 2013). The Arabidopsis genome encodes four G α genes, one canonical G α (AtGPA1) and three non-canonical extra-large Ga (XLG1, XLG2 and XLG3), a single G β gene (AGB1), three G γ genes, i.e. two typical $G\gamma$ (AGG1 and AGG2) and an atypical $G\gamma$ (AGG3), and one 7TM RGS (AtRGS1). The G γ gene duplications and evolution led to functional specialization in the plant G protein network (Chakravorty et al. 2011, Li et al. 2012, Thung et al. 2012, Trusov et al. 2008). The non-canonical $G\alpha$ proteins, XLG1, XLG2 and XLG3, have an N-terminal cysteine-rich domain and a C-terminal G α -like domain, although the G α -like domain lacks several Ga signatures required for GTP hydrolysis and G $\beta\gamma$ and RGS interactions. Fig. 1 summarizes the domain structures and the nomenclature of G protein components along with mutations discovered by forward genetics in rice.

Shoot Morphologies of Gα, RGS1 and GCR1 Mutants

Arabidopsis, rice and maize Gα mutants, gpa1, 'daikoku' dwarf1 (d1) and compact plant2 (ct2), respectively, produce shorter but wider shoot tissues (Fujisawa et al. 1999, Ullah et al. 2001, Bommert et al. 2013). The Arabidopsis gpa1 mutation confers a shortening and a widening of hypocotyls, flowers, siliques and seeds to different degrees. Fig. 2A-C presents some obvious phenotypes (e.g. leaf shape), while others (e.g. silique length) are mildly affected (Ullah et al. 2001, Ullah et al. 2003, J.G. Chen et al. 2006, Chakravorty et al. 2011). Rice and maize $G\alpha$ null alleles exhibit more severe defects; nearly all mutant shoot tissues are approximately 25-50% shorter than those of the wild-type siblings (Fujisawa et al. 1999, Bommert et al. 2013). Fig. 2D-K presents side-by-side views of the morphologies of the wild type and $G\alpha$ mutants of rice and maize. $G\alpha$ null rice DK22, one of five original rice d1 alleles (Fujisawa et al. 1999), shortens plant height by 52%, the floral bract by 25%, the seeds

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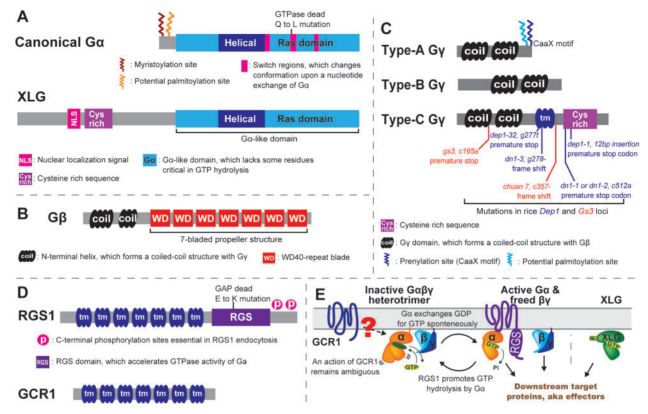


Fig. 1 Domain structures of plant G protein components. (A) Two types of G α subunits, namely canonical G α and non-canonical XLG. The G α proteins have a single $G\alpha$ domain comprising two subdomains, i.e. the Ras-homology domain and the Helical domain. Canonical $G\alpha$ has a wellconserved myristoylation site at the second glycine, and guanine nucleotide-binding motifs. Non-canonical XLG proteins have an N-terminal cysteine-rich domain, a nuclear localization signal and an unusual $G\alpha$ -like domain, which lacks some residues essential in nucleotide hydrolysis. (B) The G β subunit has N-terminal coiled-coil helices and a tryptophan-aspartic acid 40 (WD40) repeat domain. (C) Three types of G γ proteins: type-A, -B and -C G γ subunits. An N-terminal G γ domain forms a coiled-coil with the G β subunit. Type-A G γ has a well-conserved prenylation motif (CaaX motif) and a potential palmitoylation site near the C-terminus. While type-B $G\gamma$ proteins lack the prenylation motif, the rice type-B Gy protein (RGG2) is membrane associated by an unknown interaction. Type-C Gy has a transmembrane (tm) helix and a C-terminal extracellular cysteine-rich domain. Some type-C Gy proteins have a CaaX motif. Rice forward genetics identified point mutations, frameshifts and truncations in canonical Gα (RGA1, not shown) and type-C Gγ genes (DEP1 and GS3, shown in C) that confer developmental anomalies. Note that rice DEP1 and GS3 proteins vary in size (426 and 232 residues, respectively), due to a highly divergent extracellular domain. (D) Two seven transmembrane (7TM) proteins, RGS1 and GCR1. RGS1 has a 7TM region, a cytoplasmic RGS domain and C-terminal phosphorylation sites. The 7TM region has no homology to any reported GPCRs or to GCR1. GCR1 has a 7TM region, presumably having a protein fold similar to GPCRs. GCR1 is genetically uncoupled with the G protein complex in Arabidopsis development and any role for GCR1 in G protein-dependent signaling is not clear. (E) A regulatory model of the G protein complex. GDP-bound $G\alpha$ forms an inactive heterotrimer with $G\beta\gamma$ in the resting state. $G\alpha$ spontaneously exchanges GDP for GTP, releases G $\beta\gamma$ and then modulates downstream target proteins, also known as effectors. Freed G $\beta\gamma$ also modulates its own effectors. 7TM RGS1 promotes GTP hydrolysis by Gα, returning to an inactive state. An action of GCR1 on the G protein complex remains equivocal. A XLG pathway is largely unknown, except the physical and genetic association with $G\beta\gamma$. The illustrations were modified from Urano et al. (2013).

by 25% and the panicles by 50% (**Fig. 2D–K**). Other *d1* mutations are frameshifts producing premature stop codons, inframe deletions, a single residue substitution (G51E) and an epigenetic silenced allele, *epi-d1*. These alleles similarly reduce shoot growth (Ashikari et al. 1999, Miura et al. 2009, Oki et al. 2009a). The maize $G\alpha$ mutant *ct2* has a semi-dwarf stature with plant height decreased by approximately 32% and erect leaves that are approximately 31% shorter than those of the wild type (Bommert et al. 2013, Urano et al. 2015b) (**Fig. 2H**). In addition, *ct2* mutants show fasciated ears with enlarged ear tips and more rows of kernels, and thicker tassel branches, with an increased density of spikelets (**Fig. 21–K**) (Bommert et al.

2013). The $G\alpha$ mutations do not cause obvious changes in the growth rate of leaves or in flowering time (Ullah et al. 2003, Trusov et al. 2008, Urano et al. 2015b). In an opposite manner to the $G\alpha$ null mutants, ectopic expression of a constitutively active $G\alpha$, $G\alpha$ -Q222L, which mutates a glutamate (Q) residue essential in GTP hydrolysis to a leucine (L), slightly expands Arabidopsis hypocotyls under darkness (Chen et al. 2003). The findings are different under low light conditions (Okamoto et al. 2001). The equivalent Q to L mutation in the rice $G\alpha$ protein slightly enhances the longitudinal growth of shoot tissues, including internodes and seeds, by <7% (Oki et al. 2005). A 7TM negative regulator of G-proteins, RGS1, also



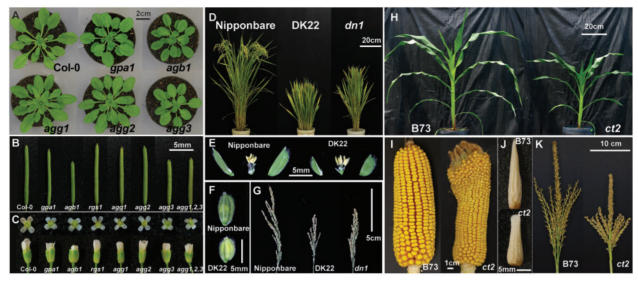


Fig. 2 Shoot morphologies of G protein mutants. (A) Rosettes of Arabidopsis seedlings grown for 40 d under short days; 8 h light at 120–130 μmol m⁻² s⁻¹ and 16 h darkness at 22°C. (B and C) Siliques and flowers of the wild type Col-0, and *gpa1-4*, *agb1-2*, *rgs1-2*, *agg1-1*, *agg2-1*, *agg3-1* and *agg1* 2 3 alleles. (D–G) Mature rice plants (D), floral architecture (E), floral bract (F) and panicles (G) of the wild type Nipponbare, the Gα-null DK22 and the Gγ (*dn1*) mutant, which lacks the cysteine-rich domain. *DN1* has two aliases: *DEP1* and *qPE9*. (H–K) Fiveweek old plants (H), mature pollinated ears (I), immature ears at approximately the V12 leaf stage (J) and tassels after anthesis (K) of wild-type B73 and Gα-null *ct2* maize. (H) is reproduced with permission from Urano et al. (2015b).

modulates shoot morphologies. Arabidopsis rgs1 null alleles, in which $G\alpha$ signal is presumably hyperactive, enhance leaf and hypocotyl outgrowths similar to the ectopic $G\alpha$ -Q222L expression (J.G. Chen et al. 2006, Chen et al. 2003), while RGS1 overexpression confers shorter hypocotyls, smaller rosettes and delayed flowering (Y. Chen et al. 2006, Johnston et al. 2007). The gpa1 rgs1 double mutant shows an epistatic interaction with the archetypical gpa1 shoot phenotypes, indicating that these two components work in the same genetic pathway (Y. Chen et al. 2006). In contrast, knockout of a putative 7TM receptor, GCR1, in the Col-0 ecotype or in the G protein mutants causes no developmental abnormality except an earlyflowering phenotype observed in an overexpression line of GCR1, suggesting no connection with the G protein complex in shoot development (Colucci et al. 2002, Chen et al. 2004, Chakraborty et al. 2015). Arabidopsis xlg3 mutants, like gpa1, displayed a shorter and wider hypocotyl (Pandey et al. 2008); however, epistasis analysis that would reveal its interaction with other G protein subunits has not been reported.

Shoot Morphologies of Gβ and Gγ Mutants

Compared with gpa1 null alleles, Arabidopsis G β null mutants, agb1, have more severe shortening of the hypocotyls, leaves, petioles, flowers, siliques and seeds (**Fig. 2A–C**), while their widths are increased to a similar level (Lease et al. 2001, Ullah et al. 2003, J.G Chen et al. 2006, Chakravorty et al. 2011). The agb1 null mutants produce more flowers (Trusov et al. 2008). The gpa1 agb1 double knockout mutants indicate an apparent epistasis of the agb1 null allele to the gpa1 null allele (J.G. Chen et al. 2006), implying that AGB1 acts downstream of GPA1, that the intact $G\alpha\beta\gamma$ complex is essential for the function or that

atypical XLGs function redundantly in the same pathway. No $G\beta$ knockout line has been isolated in rice, probably due to its embryonic lethality (Utsunomiya et al. 2012). A reduced expression of the rice $G\beta$ gene by RNA interference shortens and narrows leaf sheaths and blades (Utsunomiya et al. 2011), while the ectopic expression of $G\beta$ increases tillers and reduces leaf length (Sun et al. 2014). None of these $G\alpha$ or $G\beta$ mutations decrease cell size in shoot tissues (Ullah et al. 2001, Ullah et al. 2003, Oki et al. 2009b, Utsunomiya et al. 2011, Bommert et al. 2013); therefore, the shortened organs, caused by the $G\alpha$ or $G\beta$ mutations, are due to reduced cell proliferation (**Fig. 3A, B**).

Seed plants possess three types of $G\gamma$ subunits classified by their domain structures and lipid modification sites (Trusov et al. 2012). Type-A Gγ has a prenylation site (CaaX motif) at the C-terminus, while type-B G γ lacks this motif (Fig. 1C). Type-C Gγ has a transmembrane region and an extracellular cysteine-rich domain (Wolfenstetter et al. 2015). G β primarily co-operates with the atypical type-C G γ (e.g. Arabidopsis agg3) in shoot development. Null mutations of Arabidopsis agg3 lead to abnormal shoot morphologies, including shorter hypocotyls, siliques and seeds (Chakravorty et al. 2011, Li et al. 2012), whereas overexpression of AGG3 enlarges leaves, flowers, seeds and siliques (Li et al. 2012). Mutations in the two type-A Gγ subunits agg1 and agg2 did not lead to abnormal shoot development (Fig. 2A-C); however, AGG1 and AGG2 may still support longitudinal shoot growth, as the agg1 agg2 agg3 triple mutant shows more severe shortening of leaves, flowers and siliques than the agg3 single allele (Trusov et al. 2008, Thung et al. 2012). The agg1 agg2 agg3 triple mutant shares all the agb1 mutant shoot morphologies (Thung et al. 2012, Chakravorty et al. 2015), probably because G β is degraded in planta without G γ (Wolfenstetter et al. 2015), indicating that G γ is an



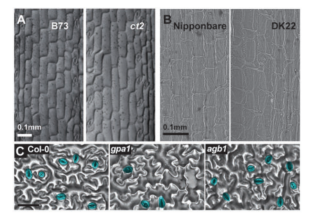


Fig. 3 Leaf epidermis of G protein mutants. (A) Epidermis of the third leaf sheath of maize B73 (wild type) and the $G\alpha$ mutant ct2. Note that the ct2 mutant has slightly longer epidermal cells. (B) Electron microscopic images of the inner epidermis of a rice floral bract. The $G\alpha$ null allele, DK22, does not change cell length. (C) Abaxial surface of Arabidopsis rosette leaves of the wild type Col-0, gpa1 and agb1. The gpa1 allele decreases while the agb1 allele increases stomatal density. Stomata are colored in cyan. Scale bar = 50 μm. The maize images were reproduced from Urano et al. (2014) with permission.

indispensable element and $G\beta$ utilizes different $G\gamma$ subtypes to sort G protein pathways.

Forward genetics studies using rice substantiate the typespecific G γ function. Rice has five G γ homologs, a type-A G γ 1 (RGG1), a type-B G γ 2 (RGG2) and three type-C G γ genes, Dense and Erect Panicle 1 (DEP1)/qPE9-1/DN1, Grain Size 3 (GS3) and Gy type-C 2 (OsGGC2) (Kato et al. 2004, Fan et al. 2006, Huang et al. 2009, Zhou et al. 2009, Taguchi-Shiobara et al. 2011, Trusov et al. 2012. Two rice quantitative trait loci (QTLs), which are associated with grain density per panicle or grain size, arise from point mutation, frameshifts or deletions of the DEP1 and GS3 genes (Fig. 1C). Similar to the type-C $G\gamma$ AGG3, DEP1 and GS3 proteins have an N-terminal G γ domain, a transmembrane region and a predicted extracellular cysteinerich domain. A premature stop codon of GS3 in the middle of the G γ domain (c165a, TGC>TGA, Fig. 1C) confers increased grain length by approximately 10%, whereas several premature terminations or frameshifts in the cysteine-rich domain (e.g. a 1 bp deletion at c357, Fig. 1C) decrease grain length (Fan et al. 2006, Takano-Kai et al. 2009, Mao et al. 2010, Takano-Kai et al. 2013). The c165a allele (gs3-3, also known as Minghui 63) is a recessive loss-of-function mutation. Suppression of the shortgrain gs3 gene (the c357- allele, gs3-4) by RNA interference expands grain length (Mao et al. 2010), suggesting that the GS3 protein gains a function by the elimination of the cysteine-rich domain. Another Gγ gene, DEP1/qPE9-1/DN1, regulates plant height, panicle erectness, and grain density and yield (Huang et al. 2009) (Fig. 2D, G). The dep1-1 allele, whose protein product lacks the entire cysteine-rich domain, increases grain quantity and primary and secondary branches per panicle, and enlarges shoot apical meristems while decreasing plant height, panicle length and grain weight (Huang et al. 2009). Other DEP1 mutations, which similarly truncate the protein, demonstrate comparable phenotypes (Zhou et al. 2009,

Taguchi-Shiobara et al. 2011, Sun et al. 2014). The dep1-1 allele is partially dominant, as ectopic expression of the truncated DEP1 protein recapitulates all the phenotypes in the near isogenic line (Sun et al. 2014), whereas dep1-32 (g277t, GGA>TGA) that expresses a G γ domain and a few residues of the transmembrane region is a recessive loss-of-function allele. These observations lead to the proposition of a model whereby the cysteine-rich domain inhibits the G β /DEP1 or G β / GS3 signals, and that eliminating part of or the entire cysteinerich domain releases the G $\beta\gamma$ dimers from this autoinhibition (Botella 2012). Rice plants overexpressing RGG1, RGG2 or GS3 are shorter compared with the parental line, although this effect has not been quantified (Mao et al. 2010, Sun et al. 2014). Further mutant analyses, including loss-of function alleles for RGG1, RGG2 or OsGGC2 genes, are necessary for understanding of the G protein network in rice development.

Meristem Activities in G Protein Mutants

G proteins are firmly established as being involved in the control mechanism for cell proliferation. The increased shoot branches of rice are related to enhanced cell proliferation or reduced determinacy of meristems. The rice Gy dep1 mutant has an enlarged inflorescence meristem (Huang et al. 2009), with increased panicle branches. Maize $G\alpha$ also regulates both the shoot apical meristem (SAM) and inflorescence meristem (IM). The maize Gα mutant ct2 has enlarged SAMs; however, their identity and organization are normal, as determined by KNOTTED1 expression analyses (Bommert et al. 2013). ct2 ear primordia have enlarged IMs, starting very early in development, leading to the initiation of extra rows of spikelet pair meristems. The tassel IMs of ct2 are also larger (Bommert et al. 2013). Abnormal meristems are similarly produced in Arabidopsis G protein mutants. While Arabidopsis gpa1 mutants display no obvious change in SAM height, the agb1 or agg1 agg2 double null alleles have approximately 40% taller meristems (Ishida et al. 2014). Both maize CT2 and Arabidopsis AGB1 function in the CLAVATA pathway, and transmit CLAVATA3 ligand-dependent signals to control meristem size, through leucine-rich repeat receptors for CLAVATA3, maize FASCIATED EAR2 or Arabidopsis Receptor-like kinase2 (Bommert et al. 2013, Ishida et al. 2014). Although these studies suggest that the G protein network co-operates with CLAVATA receptors to regulate stem cell fate, further studies are needed to understand fully the roles of G proteins in meristem regulation.

Stomatal Development in G Protein Mutants

The Arabidopsis G protein network also regulates stomata formation, most probably through control of cell proliferation, but a role in differentiation is not excluded (Fig. 3C). The Arabidopsis *gpa1* null alleles decrease stomatal density by 20–30% (Zhang et al. 2008, Nilson and Assmann 2010), while the constitutively active GPA1-Q222L mutant produces five times more stomata in the hypocotyl epidermis (Okamoto



et al. 2001) and approximately 10% more in cotyledons (Zhang et al. 2008). The rgs1 null allele similarly enhances stomatal density (Zhang et al. 2008), probably due to increasing the steady-state GPA1 activity. In contrast to the gpa1 null allele, the Arabidopsis agb1 null mutant shows slight stomatal clustering, and increased stomatal density by 25% (Zhang et al. 2008). The $G\alpha$ and $G\beta$ pathways seem to control stomatal production in cotyledons antagonistically, because the gpa1 and agb1 mutations display an additive effect on stomata formation (Zhang et al. 2008).

The role of $G\beta$ in stomatal development is coupled primarily with the typical G γ gene, AGG1. Loss-of-function alleles of agg1, but not agg2 or agg3, promoted stomatal proliferation to a level similar to agb1 (Chakravorty et al. 2015). Interestingly, the agg1 agg2 double mutant exhibited the highest stomatal density, even greater than the agb1 or agg1 agg2 agg3 triple mutant (Chakravorty et al. 2015), implying that the typical G γ subunit suppresses while the atypical $G\gamma$ subunit partially promotes stomatal development. The xlg1 xlg2 xlg3 triple knockout, but none of the xlg single null alleles, also enhances stomatal formation (Chakravorty et al. 2015). Epistasis analysis with the $\mbox{G}\beta$ or $G\gamma$ null alleles has not been tested. Insights into the underlying cellular mechanisms have come from findings that the gpa1 null mutations delay and agb1 null mutations promote asymmetric cell divisions during stomatal lineage progression (Zhang et al. 2008). Further research over successive developmental stages should elucidate how these G protein mutants alter stomatal proliferation at a molecular level.

Root Morphologies of G Protein Mutants

Arabidopsis, rice and maize $G\alpha$ null alleles decrease root growth similarly, despite their different root architectures, namely taproots in Arabidopsis vs. fibrous roots in rice and maize (Ullah et al. 2003, Izawa et al. 2010, Urano et al. 2015b). The Arabidopsis gpa1 mutant has a normal primary root length but fewer lateral roots, leading to a more compact root architecture (Ullah et al. 2003, J.G. Chen et al. 2006) (Fig. 4A), although the gpa1 effect is subtle and therefore often is overlooked with agar plate-based assays. The null alleles in rice (d1) and maize (ct2; Fig. 4B) also exhibit a slight reduction in root growth, approximately 10% shorter roots and 15% fewer seminal or crown roots compared with their wild-type sibs (Izawa et al. 2010, Urano et al. 2015b). The $G\alpha$ -null mutations probably lead to a decrease in cell proliferation at the root apical meristem, because $G\alpha$ function does not affect root cell elongation (Izawa et al. 2010). The ectopic Gα-Q222L mutation promotes primary root elongation in the opposite way due to increased cell proliferation (Chen et al. 2003).

The Arabidopsis *agb1* null mutant shows a more expanded root architecture, presumably due to increased cell proliferation and lateral root formation (Ullah et al. 2003) (**Fig. 4A**). The *agb1* phenotype is epistatic to *gpa1*, because the root architecture of the *gpa1 agb1* double mutant resembles that of the *agb1* mutant (J.G. Chen et al. 2006). AGB1 overexpression decreases lateral root formation, opposite to the loss-of-function phenotype (J.G. Chen et al. 2006). The *rgs1* null allele accelerates

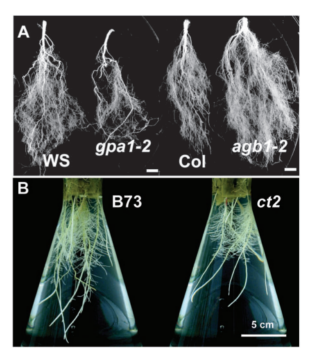


Fig. 4 Root morphologies of G protein mutants. (A) Root architecture of Arabidopsis Gα or Gβ null alleles. Scale bars = 5 mm. Note that the *gpa1-2* mutant has larger while the *agb1-2* mutant has smaller root systems. The *agb1-2* mutant and the *agg1 agg2* double mutants show increased lateral root proliferation on an agar plate. (B) Root architecture of a maize Gα mutant. The wild type B73 and Gα null *ct2* were hydroponically grown for 16 d under a daily light cycle of 16 h light at 210–220 μmol m⁻² s⁻¹ and 8 h darkness at 28°C. The Arabidopsis and maize images are adapted from Ullah et al. (2003) and Urano et al. (2015b).

primary root elongation but does not affect lateral root formation (J.G. Chen et al. 2006), whereas the gcr1 null alleles show no defect in root development or in shoot development (Pandey et al. 2006, Pandey et al. 2008), again questioning its potential involvement in G protein signaling. The xlg1 xlg2 xlg3 triple null mutant, like agb1, has a longer primary root and more lateral roots (Ding et al. 2008), although the two genotypes should be compared under the same growing conditions. However, xlg1, xlg2 or xlg3 single or double knockouts show barely changed root growth, presumably due to redundancy (Ding et al. 2008). The Arabidopsis G protein complex uses $G\gamma$ subunits spatially in shoot and root development. While the atypical AGG3 gene plays a main role in shoot development (see above), the two typical AGG1 and AGG2 genes mainly contribute to root development (Trusov et al. 2007), particularly to lateral root formation. The agg1 or agg2 mutants produce more lateral roots than Col-0, and the double mutants additively increase lateral roots to a level comparable with the agb1 allele (Trusov et al. 2007). It remains untested if this functional selectivity for $G\gamma$ subunits occurs similarly in rice and other plants.

Summary

Arabidopsis, rice and maize G protein mutants display comparable morphological anomalies, despite their distinct plant architectures. Consistent defects observed in G protein mutants are



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Morphology	
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Mutant	Shoot morphology	Root morphology	Others
Arabidopsis			
gpa1	Shorter and wider hypocotyls, leaves, seeds, and siliques with blunt tip. Opened apical hook of dark-grown hypocotyls (Ullah et al. 2001, Chen et al. 2003, Chen et al. 2006, Trusov et al. 2008, Chakravorty et al. 2011)	Reduced root mass, fewer lateral roots (Ullah et al. 2003, Chen et al. 2006)	Lower stomatal density (Zhang et al. 2008, Nilson and Assmann 2010)
xlg1 xlg2 xlg3	Shorter and wider dark-grown hypocotyls (x/g3) (Pandey et al. 2008)	More lateral roots, longer primary roots (Ding et al. 2008, Pandey et al. 2008)	Higher stomatal density (Chakravorty et al. 2015)
agb1	Shorter and wider hypocotyls, leaves, seeds, and siliques with blunt tip. Shorter mature plants, flowers, sepals and petals. Opened apical hook of dark-grown hypocotyls (Lease et al. 2001, Ullah et al. 2001, Chen et al. 2003, Chen et al. 2006a, Trusov et al. 2008, Chakravorty et al. 2011)	Increased root mass, more lateral roots, longer primary roots (Ullah et al. 2003, Chen et al. 2006, Pandey et al. 2008)	Larger shoot apical meristem (Ishida et al. 2014), higher stomatal density (Zhang et al. 2008), late flowering (Trusov et al. 2008)
gpa1 agb1	Phenocopies agb1 (Chen et al. 2006a)	Phenocopies agb1 (Chen et al. 2006, Pandey et al. 2008)	Phenocopies agb1 (Zhang et al. 2008)
agg1	Wild-type-like hypocotyls, leaves, petioles, flowers and siliques (Trusov et al. 2008)	More lateral roots (Trusov et al. 2007)	Higher stomatal density (Chakravorty et al. 2015)
agg2	Wild-type-like hypocotyls, leaves, petioles, flowers and siliques (Trusov et al. 2008)	More lateral roots (Trusov et al. 2007)	Wild-type-like stomatal density (Chakravorty et al. 2015)
agg3	Shorter and wider leaves, shorter hypocotyls (Chakaravorty et al. 2011, Thung et al. 2012),	Wild-type-like roots (Chakravorty et al. 2011)	Wild-type-like stomatal density (Chakravorty et al. 2011, Chakravorty et al. 2015)
agg1 agg2	Wild-type-like hypocotyls, leaves, petioles, flowers and siliques (Trusov et al. 2008)	More lateral roots (Trusov et al. 2007)	Higher stomatal density (Chakravorty et al. 2015)
agg1 agg2 agg3	Shorter and wider leaves, shorter siliques and flowers (Thung et al. 2012)		Higher stomatal density (Chakravorty et al. 2015)
rgs1	Longer etiolated hypocotyls, leaves and seeds (Chen et al. 2003, Chen et al. 2006a)	Longer primary roots (Chen et al. 2003)	Higher stomatal density (Zhang et al. 2008)
gcr1	Wild-type-like hypocotyls, plant height, leaves and siliques (Chen et al. 2004; Chakraborty et al. 2015)	Wild-type-like roots (Pandey et al. 2008)	
35S::GPA1		Shorter primary roots (Chen et al. 2006a)	
35S::GPA1-Q222L	Longer etiolated hypocotyls (Chen et al. 2003)	Longer primary roots (Chen et al. 2003)	Higher stomatal density (Zhang et al. 2008)
35S::RGS1	Shorter etiolated hypocotyls, smaller rosette (Chen et al. 2003, Chen et al. 2006b)	Wild-type-like root length (Chen et al. 2006b)	Late flowering (Chen et al. 2006b)
35S::GCR1			Early flowering (Colucci et al. 2002)
35S::AGB1		Shorter primary roots (Chen et al. 2006a)	Lower stomatal density (Zhang et al. 2008)
Rice			
d1 (rga1)	Shorter and wider leaves, shorter flowers, panicles and seeds. An erect panicle (Ashikari et al. 1999, Fujisawa et al. 1999, Izawa et al. 2010)	Shoter roots, fewer crown roots (Izawa et al. 2010)	



Table 1 Continued	pa		
Mutant	Shoot morphology	Root morphology	Others
rgb1 RNAi	Shorter mature plants, panicles. Shorter and narrower seeds. Brown lamina joint regions and nodes (Utsunomiya et al. 2011)		
gs3	Longer, wider and heavier seeds (gs3-3, Minghui 63), or shorter seeds (gs3-4, Chuan 7) (Fan et al. 2006, Mao et al. 2010, Takano-Kai et al. 2019, Takano-Kai et al. 2013)		
dep1/Dn1	Shorter mature plants, leaves andinflorescence internodes. More primary branches, secondary branches, and seeds per panicle (Huang et al. 2009; Taguchi-Shiobara et al. 2011, Sun et al. 2014)		Larger inflorescence meristem (Huang et al. 2009)
pActin::RGA1	Wild-type-like plant height (Sun et al. 2014)		
35S::RGA1-Q223L	Longer and heavier seeds, longer internodes (355::RGA1-Q223L in d1 back-ground) (Oki et al. 2005)		
pActin::RGG1	Shorter mature plants (Sun et al. 2014)		
pActin::RGG2	Shorter mature plants (Sun et al. 2014)		

more compact shoot architectures and altered branching patterns during the reproductive stages. The reduced organ sizes are due to lower cell proliferation activity along the longitudinal axis (Ullah et al. 2001, Ullah et al. 2003, Oki et al. 2009b, Utsunomiya et al. 2011), while changes in branching patterns are associated with enlarged meristems (Huang et al. 2009, Bommert et al. 2013, Ishida et al. 2014) and could in part be explained by control of $G\alpha$ by a master regulator of branching, as evident in the case of the regulation of maize CT2 expression by the RAMOSA1 gene (Eveland et al. 2014). The G protein complex modulates longitudinal growth potential in response to environmental factors such as light, temperature, nutrients and ions (Urano et al. 2013). This idea is supported by evidence that the maize $G\alpha$ null ct2 mutant shows effects resembling the inhibitory effect of sodium chloride on cell proliferation (Urano et al. 2014), and the rice Gγ mutant dep1 also phenocopies the growth inhibition caused by nitrogen deficiency (Sun et al. 2014). Classical plant hormone pathways including auxin, abscisic acid and gibberellin also co-ordinate with the G protein complex in various developmental processes (Urano et al. 2013). Future research should elucidate: (i) the cell type-specific function of the G protein network in cell proliferation; (ii) their co-ordination with environmental factors with regard to cell proliferation; and (iii) the regulatory systems of the G protein network in greater depth.

There are also important differences between species. For example, maize $G\alpha$ mutants have larger shoot meristems, but similar phenotypes are not seen in Arabidopsis (Bommert et al. 2013, Ishida et al. 2014). Some of these differences could be due to redundancy, as plants increased the repertoire of G protein genes during evolution, while deleting some genes in specific lineages, resulting in diversity in this signaling system. For example, Arabidopsis and its close relatives lack the type-B Gγ gene (Trusov et al. 2012), and most cereals lack the RGS gene (Urano et al. 2015a). The observed natural variation in primary structures presumably makes G protein interactions selective and signaling outputs specific. The lack of a 7TM RGS gene in cereals makes research with rice and maize of paramount importance, because no regulatory element has been identified. Experimental evidence with multiple models will lead to unexpected discoveries as well as strengthening of our current knowledge of G protein function during plant development. These hopefully will translate into improvements in crop architecture for increased harvest index.

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Disclosures

The authors have no conflicts of interest to declare.

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